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L7 19 DUP REM L5 (48 DUPLICATES REMOVED)

=> d L7 1-19 bib

L7 ANSWER 1 OF 19 BIOENG COPYRIGHT 2007 CSA on STN DUPLICATE 1
AN 2006021439 BIOENG
DN 6718654
TI Facioscapulohumeral muscular dystrophy in mice overexpressing
FRG1
AU Gabellini, Davide; D'Antona, Giuseppe; Moggio, Maurizio; Prella,
Alessandro; Zecca, Chiara; Adami, Raffaella; Angeletti, Barbara; Ciscato,
Patrizia; Pellegrino, Maria Antonietta; Bottinelli, Roberto; Green,
Michael R; Tupler, Rossella
CS Howard Hughes Medical Institute, Programs in Gene Function and Expression
and Molecular Medicine, University of Massachusetts Medical School,
Worcester, Massachusetts 01605, USA, [mailto:rossella.tupler@umassmed.edu
]
SO Nature [Nature]. Vol. 439, no. 7079, pp. 973-977. 23 Feb 2006.
Published by: Nature Publishing Group, The Macmillan Building 4 Crinan
Street London N1 9XW UK, [mailto:feedback@nature.com],
[URL:http://www.nature.com/]
ISSN: 0028-0836
DT Journal
LA English
SL English
OS CSA Neurosciences Abstracts; Genetics Abstracts

L7 ANSWER 2 OF 19 CAPLUS COPYRIGHT 2007 ACS on STN DUPLICATE 2
AN 2005:220004 CAPLUS
DN 142:291440
TI Methods for identifying activators of D4Z4 recognition complex,
YY1, HMGB2 and nucleolin for treatment of facioscapulohumeral muscular
dystrophy
IN Tupler, Rossella G.; Green, Michael R.; Gabellini, Davide
PA USA
SO U.S. Pat. Appl. Publ., 42 pp.
CODEN: USXXCO
DT Patent
LA English
FAN.CNT 1

	PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
	-----	----	-----	-----	-----
PI	US 2005054012	A1	20050310	US 2003-686491	20031014
	WO 2005037231	A2	20050428	WO 2004-US34462	20041014
	W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW				
	RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG				
PRAI	US 2002-418024P	P	20021011		
	US 2003-686491	A	20031014		

L7 ANSWER 3 OF 19 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation on STN
 AN 2005:198447 BIOSIS
 DN PREV200500190521
 TI The D4Z4 repeat-mediated pathogenesis of facioscapulohumeral
 muscular dystrophy.
 AU van der Maarel, Silvere M. [Reprint Author]; Frants, Rune R.
 CS Med CtrCtr Human and Clin Genet, Leiden Univ, NL-2333 AL, Leiden,
 Netherlands
 maarel@lumc.nl
 SO American Journal of Human Genetics, (March 2005) Vol. 76, No. 3, pp.
 375-386. print.
 CODEN: AJHGAG. ISSN: 0002-9297.
 DT Article
 General Review; (Literature Review)
 LA English
 ED Entered STN: 25 May 2005
 Last Updated on STN: 25 May 2005

L7 ANSWER 4 OF 19 MEDLINE on STN
 AN 2005218599 MEDLINE
 DN PubMed ID: 15853025
 TI First facioscapulohumeral muscular dystrophy prenatal diagnosis in a
 Bulgarian family.
 AU Buzhkov B Ts; Vuzharova R; Dimitrova V; Dimova I; Turnev I; van der Wielen
 M; van der Maarel S; Bakker B
 SO Akusherstvo i ginekologii a, (2005) Vol. 44, No. 2, pp. 30-3.
 Journal code: 0370455. ISSN: 0324-0959.
 CY Bulgaria
 DT (CASE REPORTS)
 Journal; Article; (JOURNAL ARTICLE)
 LA Bulgarian
 FS Priority Journals
 EM 200505
 ED Entered STN: 29 Apr 2005
 Last Updated on STN: 27 May 2005
 Entered Medline: 26 May 2005

L7 ANSWER 5 OF 19 USPATFULL on STN DUPLICATE 3
 AN 2004:50778 USPATFULL
 TI Gene expression in bladder tumors
 IN Orntoft, Torben F., Aabyhoj, DENMARK
 PI US 2004038207 A1 20040226
 US 6936417 B2 20050830
 AI US 2001-951968 A1 20010914 (9)
 RLI Division of Ser. No. US 2000-510643, filed on 22 Feb 2000, UNKNOWN
 DT Utility
 FS APPLICATION
 LREP BANNER & WITCOFF, 1001 G STREET N W, SUITE 1100, WASHINGTON, DC, 20001
 CLMN Number of Claims: 26
 ECL Exemplary Claim: 1
 DRWN 15 Drawing Page(s)
 LN.CNT 28561
 CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L7 ANSWER 6 OF 19 SCISEARCH COPYRIGHT (c) 2007 The Thomson Corporation on
 STN
 AN 2004:237731 SCISEARCH
 GA The Genuine Article (R) Number: 800JT
 TI Molecular basis of facioscapulohumeral muscular dystrophy
 AU Tupler R (Reprint); Gabellini D
 CS Univ Massachusetts, Sch Med, Program Gene Funct & Express, Lazare Med Res
 Bldg, 364 Plantat St, Rm 660, Worcester, MA 01605 USA (Reprint); Univ
 Massachusetts, Sch Med, Program Gene Funct & Express, Worcester, MA 01605
 USA; Univ Pavia, I-27100 Pavia, Italy

CYA USA; Italy
 SO CELLULAR AND MOLECULAR LIFE SCIENCES, (MAR 2004) Vol. 61, No. 5, pp. 557-566.
 ISSN: 1420-682X.
 PB BIRKHAUSER VERLAG AG, VIADUKSTRASSE 40-44, PO BOX 133, CH-4010 BASEL, SWITZERLAND.
 DT General Review; Journal
 LA English
 REC Reference Count: 70
 ED Entered STN: 19 Mar 2004
 Last Updated on STN: 19 Mar 2004
 ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

L7 ANSWER 7 OF 19 MEDLINE on STN
 AN 2004534031 MEDLINE
 DN PubMed ID: 15504910
 TI The 4q subtelomere harboring the FSHD locus is specifically anchored with peripheral heterochromatin unlike most human telomeres.
 AU Tam Rose; Smith Kelly P; Lawrence Jeanne B
 CS Department of Cell Biology, University of Massachusetts Medical School, Worcester, MA 01655, USA.
 NC GM 68138 (NIGMS)
 SO The Journal of cell biology, (2004 Oct 25) Vol. 167, No. 2, pp. 269-79.
 Journal code: 0375356. ISSN: 0021-9525.
 CY United States
 DT Journal; Article; (JOURNAL ARTICLE)
 LA English
 FS Priority Journals
 EM 200412
 ED Entered STN: 27 Oct 2004
 Last Updated on STN: 19 Dec 2004
 Entered Medline: 6 Dec 2004

L7 ANSWER 8 OF 19 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation on STN DUPLICATE 4
 AN 2004:9067 BIOSIS
 DN PREV200400000121
 TI Testing the position-effect variegation hypothesis for facioscapulohumeral muscular dystrophy by analysis of histone modification and gene expression in subtelomeric 4q.
 AU Jiang, Guanchao; Yang, Fan; van Overveld, Petra G. M.; Vedanarayanan, Vettaikorumakankav; van der Maarel, Silvere; Ehrlich, Melanie [Reprint Author]
 CS Human Genetics Program and Department of Biochemistry, Tulane Medical School, New Orleans, LA, 70112, USA
 ehrlich@tulane.edu
 SO Human Molecular Genetics, (15 November 2003) Vol. 12, No. 22, pp. 2909-2921. print.
 ISSN: 0964-6906 (ISSN print).
 DT Article
 LA English
 ED Entered STN: 17 Dec 2003
 Last Updated on STN: 17 Dec 2003

L7 ANSWER 9 OF 19 MEDLINE on STN
 AN 2003342372 MEDLINE
 DN PubMed ID: 12874395
 TI D4F104S1 deletion in facioscapulohumeral muscular dystrophy: phenotype, size, and detection.
 AU Lemmers R J L F; Osborn M; Haaf T; Rogers M; Frants R R; Padberg G W; Cooper D N; van der Maarel S M; Upadhyaya M
 CS Department of Human Genetics, Center for Human and Clinical Genetics, Leiden, The Netherlands.
 SO Neurology, (2003 Jul 22) Vol. 61, No. 2, pp. 178-83.

Journal code: 0401060. E-ISSN: 1526-632X.

CY United States
 DT Journal; Article; (JOURNAL ARTICLE)
 LA English
 FS Abridged Index Medicus Journals; Priority Journals
 EM 200404
 ED Entered STN: 23 Jul 2003
 Last Updated on STN: 28 Apr 2004
 Entered Medline: 27 Apr 2004

L7 ANSWER 10 OF 19 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation on
 STN
 AN 2003:153603 BIOSIS
 DN PREV200300153603
 TI Increasing D4Z4 repeat copy number compromises C2C12 myoblast
 differentiation.
 AU Yip, Darren J.; Picketts, David J. [Reprint Author]
 CS Molecular Medicine Program, Ottawa Health Research Institute, 501 Smyth
 Road, Ottawa, ON, K1H 8L6, Canada
 dpicketts@ohri.ca
 SO FEBS Letters, (27 February 2003) Vol. 537, No. 1-3, pp. 133-138. print.
 CODEN: FEBLAL. ISSN: 0014-5793.
 DT Article
 LA English
 ED Entered STN: 26 Mar 2003
 Last Updated on STN: 26 Mar 2003

L7 ANSWER 11 OF 19 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation on
 STN DUPLICATE 5
 AN 1999:124876 BIOSIS
 DN PREV199900124876
 TI Recent amplification of the human FRG1 gene during primate
 evolution.
 AU Grewal, Prabhjit K.; Van Geel, Michel; Frants, Rune R.; De Jong, Pieter;
 Hewitt, Jane E. [Reprint author]
 CS Div. Genet., Queen's Med. Cent., Nottingham Univ., Nottingham NG7 2UH, UK
 SO Gene (Amsterdam), (Feb. 4, 1999) Vol. 227, No. 1, pp. 79-88. print.
 CODEN: GENED6. ISSN: 0378-1119.
 DT Article
 LA English
 ED Entered STN: 17 Mar 1999
 Last Updated on STN: 17 Mar 1999

L7 ANSWER 12 OF 19 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation on
 STN DUPLICATE 6
 AN 2000:2314 BIOSIS
 DN PREV200000002314
 TI The FSHD region on human chromosome 4q35 contains potential
 coding regions among pseudogenes and a high density of repeat elements.
 AU van Geel, M.; Heather, L. J.; Lyle, R.; Hewitt, J. E.; Frants, R. R.; de
 Jong, P. J. [Reprint author]
 CS Department of Cancer Genetics, Roswell Park Cancer Institute, Elm and
 Carlton Streets, Buffalo, NY, 14263, USA
 SO Genomics, (Oct. 1, 1999) Vol. 61, No. 1, pp. 55-65. print.
 CODEN: GNMCEP. ISSN: 0888-7543.
 DT Article
 LA English
 OS Genbank-AF146191; EMBL-AF146191; Genbank-U85056; EMBL-U85056;
 Genbank-U89471; EMBL-U89471
 ED Entered STN: 23 Dec 1999
 Last Updated on STN: 31 Dec 2001

L7 ANSWER 13 OF 19 BIOSIS COPYRIGHT (c) 2007 The Thomson Corporation on
 STN DUPLICATE 7

AN 1997:319965 BIOSIS
 DN PREV199799610453
 TI The mouse homolog of FRG1, a candidate gene for FSHD,
 maps proximal to the myodystrophy mutation on chromosome 8.
 AU Grewal, Prabhjit K.; Van Deutekom, Judith C. T.; Mills, Kate A.; Lemmers,
 Richard J. L. F.; Mathews, Kathy D.; Frants, Rune R.; Hewitt, Jane E.
 [Reprint author]
 CS Sch. Biol. Sci., Univ. Manchester, 3.239 Stopford Build., Oxford Road,
 Manchester M13 9PT, UK
 SO Mammalian Genome, (1997) Vol. 8, No. 6, pp. 394-398.
 CODEN: MAMGEC. ISSN: 0938-8990.
 DT Article
 LA English
 ED Entered STN: 26 Jul 1997
 Last Updated on STN: 26 Jul 1997

L7 ANSWER 14 OF 19 CAPLUS COPYRIGHT 2007 ACS on STN DUPLICATE 8
 AN 1997:74740 CAPLUS
 DN 126:127617
 TI Localization of the cell death genes CPP32 and Mch-2 to human chromosome
 4q
 AU Nasir, J.; Theilmann, J. L.; Chopra, V.; Jones, A. M.; Walker, D.; Rasper,
 D. M.; Vaillancourt, J. P.; Hewitt, J. E.; Nicholson, D. W.; Hayden, M. R.
 CS Department of Medical Genetics and Centre for Molecular Medicine &
 Therapeutics (CMMT), University of British Columbia, Vancouver, BC, V6T
 1Z4, Can.
 SO Mammalian Genome (1997), 8(1), 56-59
 CODEN: MAMGEC; ISSN: 0938-8990
 PB Springer
 DT Journal
 LA English
 RE.CNT 28 THERE ARE 28 CITED REFERENCES AVAILABLE FOR THIS RECORD
 ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 15 OF 19 BIOTECHNO COPYRIGHT 2007 Elsevier Science B.V. on STN
 DUPLICATE
 AN 1997:27122411 BIOTECHNO
 TI Molecular genetics of facioscapulohumeral muscular dystrophy. (FSHD)
 AU Fisher J.; Upadhyaya M.
 CS M. Upadhyaya, Institute of Medical Genetics, University of Wales, College
 of Medicine, Cardiff CF4 4XN, United Kingdom.
 SO Neuromuscular Disorders, (1997), 7/1 (55-62), 69 reference(s)
 CODEN: NEDIEC ISSN: 0960-8966.
 PUI S0960896696004002
 DT Journal; Article
 CY United Kingdom
 LA English
 SL English

L7 ANSWER 16 OF 19 DGENE COPYRIGHT 2007 The Thomson Corp on STN
 AN ADY54205 DNA DGENE
 TI Identifying candidate therapeutic compounds for treating
 facioscapulohumeral muscular dystrophy, by contacting D4Z4
 binding element with test compound, and determining interaction between
 test compound and D4Z4 binding element.
 IN Tupler R G; Green M R; Gabellini D
 PA (TUPL-I) TUPLER R G.
 (GREE-I) GREEN M R.
 (GABE-I) GABELLINI D.
 PI US 2005054012 AI 20050310 42
 AI US 2003-686491 20031014
 PRAI US 2002-418024P 20021011
 DT Patent

LA English
 OS 2005-221928 [23]
 DESC Human FRG1 quantitative PCR primer FRG1-lrb.

L7 ANSWER 17 OF 19 DGENE COPYRIGHT 2007 The Thomson Corp on STN
 AN ADY54204 DNA DGENE
 TI Identifying candidate therapeutic compounds for treating facioscapulohumeral muscular dystrophy, by contacting D4Z4 binding element with test compound, and determining interaction between test compound and D4Z4 binding element.
 IN Tupler R G; Green M R; Gabellini D
 PA (TUPL-I) TUPLER R G.
 (GREE-I) GREEN M R.
 (GABE-I) GABELLINI D.
 PI US 2005054012 A1 20050310 42
 AI US 2003-686491 20031014
 PRAI US 2002-418024P 20021011
 DT Patent
 LA English
 OS 2005-221928 [23]
 DESC Human FRG1 quantitative PCR primer FRG1-lf.

L7 ANSWER 18 OF 19 DGENE COPYRIGHT 2007 The Thomson Corp on STN
 AN ADY54207 DNA DGENE
 TI Identifying candidate therapeutic compounds for treating facioscapulohumeral muscular dystrophy, by contacting D4Z4 binding element with test compound, and determining interaction between test compound and D4Z4 binding element.
 IN Tupler R G; Green M R; Gabellini D
 PA (TUPL-I) TUPLER R G.
 (GREE-I) GREEN M R.
 (GABE-I) GABELLINI D.
 PI US 2005054012 A1 20050310 42
 AI US 2003-686491 20031014
 PRAI US 2002-418024P 20021011
 DT Patent
 LA English
 OS 2005-221928 [23]
 DESC Human FRG1 RT-PCR primer FRG1-lrb.

L7 ANSWER 19 OF 19 DGENE COPYRIGHT 2007 The Thomson Corp on STN
 AN ADY54206 DNA DGENE
 TI Identifying candidate therapeutic compounds for treating facioscapulohumeral muscular dystrophy, by contacting D4Z4 binding element with test compound, and determining interaction between test compound and D4Z4 binding element.
 IN Tupler R G; Green M R; Gabellini D
 PA (TUPL-I) TUPLER R G.
 (GREE-I) GREEN M R.
 (GABE-I) GABELLINI D.
 PI US 2005054012 A1 20050310 42
 AI US 2003-686491 20031014
 PRAI US 2002-418024P 20021011
 DT Patent
 LA English
 OS 2005-221928 [23]
 DESC Human FRG1 RT-PCR primer FRG1-lf.

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